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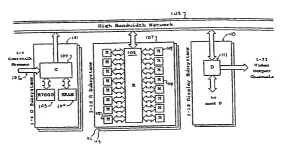
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(54) Title: NEUROTROPHIC FACTOR RECEPTOR



(57) Abstract: There is disclosed a mammalian receptor protein designated GFRc-4 and the nucleic acid sequence encoding therefor. Also provided is an expression vector comprising the nucleic acid sequence and host cells transformed or transfected with the
vector. Host cells expressing the receptor may be used to identify agonists or antagonists in relation to the receptor.

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NEUROTROPHIC FACTOR RECEPTOR

The present invention is concerned with cloning and expression of a novel mammalian receptor protein, designated herein GFR α -4 and in particular with an isolated nucleic acid sequence encoding the GFR α -4 protein, an expression vector comprising said nucleic acid sequence, a host cell transformed or transfected with said vector, isolated GFR α 4 protein, compounds which act as agonists or antagonists in relation to GFR α -4 and methods of identifying them, together with pharmaceutical compositions comprising the isolated nucleic acid, the receptor protein or said agonist or antagonist.

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Neurotrophic growth factors are involved in neuronal differentiation, development and maintenance. These proteins can prevent degeneration and promote survival of different types of neuronal cells and are thus potential therapeutic agents for neurodegenerative diseases. Glial cell-line derived neurotrophic factor (GDNF) was the first member of a growing subfamily of neurotrophic factors structurally distinct from the neurotrophins. GDNF is a distantly related member of the transforming growth factor β (TGF- β) superfamily of growth factors, characterized by a specific pattern of seven highly conserved cysteine residues within the amino acid sequence (Kingsley, 1994). GDNF was originally purified using an assay based on its ability to maintain the survival and function of embryonic ventral midbrain dopaminergic neurons in vitro (Lin et al., 1993). Other neuronal cell types in the central (CNS) or peripheral nervous systems (PNS) have been shown to be responsive to the survival effects of GDNF (Henderson et al., 1994, Buj-Bello et al., 1995, Mount et al., 1995, Oppenheim et al., 1995). GDNF is produced by cells in an inactive

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proform, which is cleaved specifically at a RXXR recognition site to produce active GDNF (Lin et al., 1993). In view of its effects on dopaminergic neurons, clinical trials have evaluated GDNF as a possible treatment for Parkinson's disease, a common neurodegenerative disorder characterized by the loss of a high percentage (up to 70 %) of dopaminergic cells in the substantia nigra of the brain. Exogenous administration of GDNF has potent protective effects in animal models of Parkinson's disease (Henderson et 10 al., 1994, Beck et al., 1995, Tomac et al., 1995, Yan et al., 1995, Gash et al., 1996, Choi-Lundberg et al., 1997, Bilang-Bleuel et al., 1997, Mandel et al., 1997).

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Recently, three new members of the GDNF family of neurotrophic factors have been discovered. Neurturin (NTN) was purified from conditioned medium from Chinese hamster ovary (CHO) cells using an assay based on the ability to promote the survival of sympathetic 20 neurons in culture (Kotzbauer et al., 1996). The mature neurturin protein is 57% similar to mature GDNF. Persephin (PSP) was discovered by degenerate primer PCR using genomic DNA. The mature protein, like mature GDNF, promotes the survival of ventral midbrain 25 dopaminergic neurons and of motor neurons in culture (Milbrandt et al., 1998). The similarity of the mature persephin protein with mature GDNF and neurturin is \approx 50 %. Very recently, a fourth member has been cloned using genomic DNA information in the public EMBL 30 database and has been named Enovin (EVN) (Masure et al., 1999) or Artemin (ARTN) (Baloh et al., 1998b). This factor is \pm 57 % similar to NTN and PSP and acts primarily on peripheral neurons.

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All four GDNF family members require a heterodimeric receptor complex in order to carry out downstream

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intracellular signal transduction. GDNF binds to the GDNF family receptor alpha 1 (GFR α -1; also termed $\mbox{GDNFR}\alpha$, RETL1 or TrnR1; $\mbox{GFR}\alpha$ Nomenclature Committee, 1997) subunit, a glycosyl phosphatidyl inositol (GPI)anchored membrane protein (Jing et al., 1996, Treanor 5 et al., 1996, Sanicola et al., 1997). The GDNF/GFR α -1 complex subsequently binds to and activates the cRET proto-oncogene, a membrane bound tyrosine kinase (Durbec et al., 1996, Trupp et al., 1996), resulting in phosphorylation of tyrosine residues in cRET and 10 subsequent activation of downstream signal transduction pathways (Worby et al., 1996). GFR α -2 (also termed RETL2, NTNR- α , GDNFR- β or TrnR2), which is similar to GFR α -1, has been identified by a number of different groups (Baloh et al., 1997, Sanicola et 15 al., 1997, Klein et al., 1997, Buj-Bello et al., 1997, Suvanto et al., 1997). The human GFR α -1 and GFR α -2 receptor subunits are 49% identical and 63% similar by protein sequence with 30 of the 31 cysteine residues conserved. Both receptors contain a hydrophobic domain 20 at their carboxy-termini involved in GPI anchoring to the membrane. GFR α -1 and GFR α -2 are widely expressed in almost all tissues and expression may be developmentally regulated (Sanicola et al., 1997, Widenfalk et al., 1997). 25

GFR α -1 is the preferred receptor for GDNF, whereas GFR α -2 preferentially binds neurturin (Jing et al., 1996, Treanor et al., 1996, Klein et al., 1997). It is also clear, however, that there is some cross-talk between these growth factors and receptors as GDNF can bind to GFR α -2 in the presence of cRET (Sanicola et al., 1997) and neurturin can bind to GFR α -1 with low affinity (Klein et al., 1997). GDNF and neurturin are thus part of a neurotrophic signalling system whereby different ligand-binding subunits (GFR α -1 and GFR α -2) can interact with the same tyrosine kinase subunit

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(cRET).

Recently, a third member of the $GFR\alpha$ family of coreceptors, $GFR\alpha-3$, has been described (Jing et al., 1997, Masure et al., 1998, Worby et al., 1998, 5 Naveilhan et al., 1998, Baloh et al., 1998a). This receptor's amino acid sequence is 35% identical to both GFR α -1 and GFR α -2. GFR α -3 is not expressed in the developing or adult CNS, but is highly expressed in several developing and adult sensory and sympathetic 10 ganglia of the PNS (Widenfalk et al., 1998, Naveilhan et al., 1998, Baloh et al., 1998a). GFR α -3 has been shown to be the preferred coreceptor for Enovin/artemin and also signals via cRET (Masure et al., 1999, Baloh et al., 1998b). Crosstalk between 15 EVN/ARTN and GFR α -1 seems also possible, at least in vitro.

A fourth member of the GFR α family has been identified in chicken (Thompson et al., 1998) and has been shown to mediate signalling of persephin via cRET (Enokido et al., 1998). A functional mammalian homologue encoding a mammalian persephin receptor has yet to be discovered.

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The present inventors have surprisingly identified a further novel mammalian receptor of the GDNF family designated herein as $GFR\alpha-4$. The DNA sequence has been cloned and a number of splice variants encoding the receptor have also been identified.

Accordingly, there is provided by the present invention an isolated or substantially pure form of a nucleic acid encoding a mammalian GDNF family receptor $\alpha-4$ designated GFR $\alpha-4$. The nucleic acid molecule is preferably from rat, mouse or human. Preferably, the receptor encoded by said nucleic acid molecule

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comprises the amino acid sequence illustrated in Sequence ID No's. 8 or 9 or encoding a functional equivalent, derivative or bioprecursor of said receptor.

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Although initially, in view of the fact that only 4 members of the GFRas were known, the new receptor has previously been called $GFR\alpha-5$. However, it has now termed α -4 in order to comply with the existing nomenclature for $GFR\alpha$ family members and to indicate that the GFR α receptor of the present invention is the mammalian orthologue of the chicken $\ensuremath{\mathsf{GFR}\alpha\mbox{-4}}$ receptor.

Thus, the present invention relates to a nucleic acid molecule encoding a mammalian GDNF family receptor α -4 (GFR α -4) or an immunologically and/or biologically active fragment thereof, which comprises a nucleotide sequence selected from the group consisting of

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(a) nucleotide sequences encoding the polypeptide comprising the amino acid sequence depicted in SEQ ID NO: 8 or 9; nucleotide sequences comprising the coding

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(b)

sequence as depicted in SEQ ID NO: 5 or 6; nucleotide sequences encoding a polypeptide (c) derived from the polypeptide encoded by a nucleotide sequence of (a) or (b) by way of substitution, deletion and/or addition of one or several amino acids of the amino acid sequence encoded by the nucleotide sequence

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nucleotide sequences the complementary (d) strand of which hybridizes with a nucleotide sequence of any one of (a) to (c);

of (a) or (b);

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nucleotide sequences encoding a polypeptide (e) the amino acid sequence of which has an identity of 30% or more to the amino acid sequence of the polypeptide encoded by a

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nucleotide sequence of any one of (a) to (d);

- (f) nucleotide sequences encoding a polypeptide capable of binding persephin comprising a fragment or an epitope-bearing portion of a polypeptide encoded by a nucleotide sequence of any one of (a) to (e);
 - (g) nucleotide sequences comprising at least 15 consecutive nucleotides of a nucleotide sequence of any one of (a) to (f) and encoding a polypeptide capable of binding persephin; and
 - (h) nucleotide sequences comprising a nucleotide sequence which is degenerated as a result of the genetic code to a nucleotide sequence of any of (a) to (g).

Advantageously, the nucleic acid molecule according to the invention may be used for expression of said $\mbox{GFR}\alpha\mbox{-4}$ protein in, for example, a host cell or the 20 like, using an appropriate expression vector. Preferably, the nucleic acid molecule is a DNA molecule, and even more preferably a cDNA molecule having a sequence as illustrated in any of Sequence ID No's. 5 to 7 or the complement thereof. 25 Alternatively, the nucleic acid molecule is capable of hybridising to the sequences of the invention under conditions of high stringency or to the complement thereof. Stringency of hybridisation as used herein refers to conditions under which polynucleic acids are 30 stable. The stability of hybrids is reflected in the melting temperature (Tm) of the hybrids. Tm can be approximated by the formula:

wherein 1 is the length of the hybrids in nucleotides. Tm decreases approximately by 1-1.5°C with every 1% decrease in sequence homology.

5 The nucleic acid capable of hybridising to nucleic acid molecules according to the invention will generally be at least 70%, preferably at least 80 or 90% and more preferably at least 95% homologous to the nucleotide sequences according to the invention.

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Advantageously, the antisense molecule may be used as a probe or as a medicament or in a pharmaceutical composition together with a pharmaceutically acceptable carrier, diluent or excipient.

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According to a second aspect of the invention, there is provided a DNA expression vector comprising the DNA molecule according to the invention. This vector may, advantageously, be used to transform or transfect a host cell to achieve expression of $GFR\alpha-4$ according to the invention. Preferably, the DNA is included in a plasmid, for subsequent transfection or transformation of the host cell.

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An expression vector according to the invention includes a vector having a nucleic acid according to the invention operably linked to regulatory sequences, such as promoter regions, that are capable of effecting expression of said DNA fragments. The term "operably linked" refers to a juxta position wherein the components described are in a relationship permitting them to function in their intended manner. Such vectors may be transformed into a suitable host cell to provide for expression of a polypeptide according to the invention. Thus, in a further aspect, the invention provides a process for preparing receptors according to the invention which comprises

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cultivating a host cell, transformed or transfected with an expression vector as described above under conditions to provide for expression by the vector of a coding sequence encoding the receptors, and recovering the expressed receptors.

The vectors may be, for example, plasmid, virus or phage vectors provided with an origin of replication, optionally a promoter for the expression of said nucleotide and optionally a regulator of the promoter. The vectors may contain one or more selectable markers, such as, for example, ampicillin resistance.

Regulatory elements required for expression include promoter sequences to bind RNA polymerase and 15 transcription initiation sequences for ribosome binding. For example, a bacterial expression vector may include a promoter such as the lac promoter and for transcription initiation in the Shine-Dalgarno sequence and the start codon AUG. Similarly, a 20 eukaryotic expression vector may include a heterologous or homologous promoter for RNA polymerase II, a downstream polyadenylation signal, the start codon AUG, and a termination codon for detachment of the ribosome. Such vectors may be obtained 25 commercially or assembled from the sequences described by methods well known in the art.

Nucleic acid molecules according to the invention may be inserted into the vectors described in an antisense orientation in order to provide for the production of antisense RNA. Antisense RNA or other antisense nucleic acids may be produced by synthetic means.

35 In accordance with the present invention, a defined nucleic acid includes not only the identical nucleic acid but also any amino base variations including, in WO 01/02557

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particular, substitutions in bases which result in a synonymous codon (a different codon specifying the same amino acid residue) due to the degenerate code in conservative amino acid substitutions. The term "nucleic acid sequence" also includes the complementary sequence to any single stranded sequence given regarding base variations.

The present invention also advantageously provides nucleic acid sequences of at least approximately 10 10 contiguous nucleotides of a nucleic acid according to the invention and preferably from 10 to 50 nucleotides. These sequences may, advantageously, be used as probes or primers to initiate replication, or the like. Such nucleic acid sequences may be produced 15 according to techniques well known in the art, such as, by recombinant or synthetic means. They may also be used in diagnostic kits or the like for detecting the presence of a nucleic acid according to the invention. These tests generally comprise contacting 20 the probe with the sample under hybridising conditions and detecting for the presence of any duplex or triplex formation between the probe and any nucleic acid in the sample.

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According to the present invention these probes may be anchored to a solid support. Preferably, they are present on an array so that multiple probes can simultaneously hybridize to a single biological sample. The probes can be spotted onto the array or synthesised in situ on the array. (See Lockhart et al., Nature Biotechnology, vol. 14, December 1996 "Expression monitoring by hybridisation into high density oligonucleotide arrays". A single array can contain more than 100, 500 or even 1,000 different probes in discrete locations.

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The nucleic acid sequences, according to the invention may be produced using such recombinant or synthetic means, such as, for example, using PCR cloning mechanisms which generally involve making a pair of primers, which may be from approximately 10 to 50 5 nucleotides to a region of the gene which is desired to be cloned, bringing the primers into contact with mRNA, cDNA, or genomic DNA from a human cell, performing a polymerase chain reaction under conditions which bring about amplification of the 10 desired region, isolating the amplified region or fragment and recovering the amplified DNA. Generally, such techniques as defined herein are well known in the art, such as described in Sambrook et al (Molecular Cloning: a Laboratory Manual, 1989). 15

The nucleic acids or oligonucleotides according to the invention may carry a revealing label. Suitable labels include radioisotopes such as ³²P or ³⁵S, enzyme labels or other protein labels such as biotin or fluorescent markers. Such labels may be added to the nucleic acids or oligonucleotides of the invention and may be detected using known techniques per se.

25 The present invention also comprises within its scope proteins or polypeptides encoded by the nucleic acid molecules according to the invention or a functional equivalent, derivative or bioprecursor thereof.

Preferably, the protein comprises the amino acid sequence of Sequence ID No's. 8 and 9.

A "functional equivalent" as defined herein should be taken to mean a receptor that exhibits the same properties and functionality associated with the ${\sf GFR}\alpha-4$ receptor according to the invention. A "derivative" should be taken to mean a polypeptide or protein in which certain amino acids may have been

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altered or deleted or replaced and which polypeptide or protein retains biological activity of said $GFR\alpha-4$ receptor and/or which can cross react with antibodies raised using a receptor according to the invention as the challenging antigen.

Encompassed within the scope of the invention are hybrid and modified forms of the $GFR\alpha-4$ receptor according to the invention including fusion proteins and fragments. The hybrid and modified forms include, for example, when certain amino acids have been subjected to some modification or replacement, such as for example, by point mutation and yet which results in a protein which possesses the same receptor specificity as the $GFR\alpha-4$ receptor of the invention.

In this context it is understood that the terms biological activity, receptor specificity and functional receptor (fragment) preferably include the ability to bind persephin, preferably specifically and that the $GFR\alpha-4$ of the invention has no or substantially no binding activity on GDNF, NTN, and/or EVN/ART. The biological activity, receptor specificity and/or functionality, for example binding activity of the $GFR\alpha$ -4 of the invention, variants, derivatives and fragments thereof can be determined according methods well known in the art, preferably as described in the appended examples. Preferably, the $K_{\scriptscriptstyle D}$ of the $GFR\alpha\text{--}4$ of the invention for persephin is 1 to 10 x 10^{-9} , particularly preferred 5.9 \pm 2.8 x $10^{-9}\,\mathrm{when}$ determined , in accordance with the examples described below; see the description to Table 6.

The protein according to the invention should be taken to include all possible amino acid variants encoded by the nucleic acid molecule according to the invention

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including a polypeptide encoded by said molecule and having conservative amino acid changes. Proteins or polypeptides according to the invention further include variants of such sequences, including naturally occurring allelic variants which are substantially homologous to said proteins or polypeptides. In this context, substantial homology is regarded as a sequence which has at least 70%, and preferably 80 or 90% amino acid homology with the proteins or polypeptides encoded by the nucleic acid molecules according to the invention.

Substantial homology should be taken to mean that the nucleotide and amino acid sequences of the $\ensuremath{\mathsf{GFR}\alpha}\xspace-4$ of the invention display a certain degree of sequence 15 identity. Preferably they share an identity of at least 30 %, preferably 40 %, more preferably 50 %, still more preferably 60 %, most preferably 70%, and particularly an identity of at least 80 %, preferably more than 90 % and still more preferably more than 95 $\,$ 20 % is desired with respect to the nucleotide or amino acid sequences depicted in Seq. ID Nos. 5 to 9, respectively. A preferred method for determining the best overall match between a query sequence (a sequence of the present invention) and a subject 25 sequence, also referred to as a global sequence alignment, can be determined using, for example, the FASTDB computer program based on the algorithm of Brutlag et al. (Comp. App. Biosci. 6 (1990), 237-245.) In a sequence alignment the query and subject 30 sequences are both DNA sequences. An RNA sequence can be compared by converting U's to T's. The result of said global sequence alignment is in percent identity. Further programs that can be used in order to determine homology/identity are described below and in 35 the examples. The sequences that are homologous to the sequences described above are, for example, variations WO 01/02557 PCT/EP00/04918

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of said sequences which represent modifications having the same biological function, in particular encoding proteins with the same or substantially the same receptor specificity, i.e. binding specificity. They may be naturally occurring variations, such as 5 sequences from other mammals, or mutations. These mutations may occur naturally or may be obtained by mutagenesis techniques. The allelic variations may be naturally occurring allelic variants as well as synthetically produced or genetically engineered 10 variants. In a preferred embodiment the sequences are derived from mouse, more preferable from human. These sequences can also be retrieved from existing databases with nucleotide sequences of yet unknown function. For example, a BLAST search on the EMBL 15 database using the identified rat $GFR\alpha-4$ sequences as query sequence yielded a genomic mouse sequence (accession no. AF155960) and a genomic human sequence (accession no. AC017113; containing contigs. derived from human genomic DNA) with parts almost identical 20 with parts of the rat $GFR\alpha-4$ sequence (exons 2, 3 and 4). These nucleotides sequences are also encompassed in the present invention.

A further aspect of the invention comprises the host cell itself transformed with the DNA expression vector described herein, which host cell preferably comprises a eukaryotic cell, which may be for example, a mammalian cell, an insect cell or yeast cell or the like. In one embodiment the cell comprises a human embryonic kidney cell and preferably a cell of the HEK293 cell line. Alternatively, the cell may comprise NIH/3T3 mouse fibroblasts or Chinese hamster ovary (CHO) cells or COS-7 cells.

Further provided by the present invention is a transgenic cell, tissue or organism comprising a

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transgene capable of expressing $GFR\alpha-4$ according to the invention, or of expressing a functional equivalent, derivative or bioprecursor of said receptor. The term "transgene capable of expression" as used herein means a suitable nucleic acid sequence 5 which leads to expression of a human receptor having the same function and/or activity as $\mbox{GFR}\alpha\mbox{-4.}$ The transgene may include, for example, genomic nucleic acid isolated from rat cells or synthetic nucleic acid, including cDNA, integrated into the genome or in 10 an extra chromosomal state. Preferably, the transgene comprises the nucleic acid sequence encoding $\mbox{GFR}\alpha\mbox{-}4$ according to the invention or a functional fragment of said nucleic acid. A functional fragment of said nucleic acid should be taken to mean a fragment of the 15 gene or cDNA encoding $GFR\alpha-4$ receptor or a functional equivalent or bioprecursor of said $GFR\alpha-4$ which fragment is capable of being expressed to produce a functional receptor protein. For example, the gene may comprise deletions of mutations but may still 20 encode a functional receptor.

Further provided by the present invention is an isolated or purified $GFR\alpha-4$ protein having the amino acid sequence illustrated in Sequence ID No. 2 or a functional fragment or bioprecursor of said receptor or alternatively a $GFR\alpha-4$ protein expressed by the transgenic cell, tissue or organism according to the invention. Also provided by the invention are membrane preparations from cells expressing $GFR\alpha-4$.

The present invention is further directed to inhibiting GFRα-4 in vivo by the use of antisense technology. Antisense technology can be used to control gene expression through triple-helix formation or antisense DNA or RNA, both of which methods are based on binding of a polynucleotide to DNA or RNA.

For example, the 5' coding portion of the mature protein sequence, which encodes for the protein of the present invention, is used to design an antisense RITA oligonucleotide of from 10 to 40 base pairs in length.

5 A DNA oligonucleotide is designed to be complementary to a region of the gene involved in transcription (triple-helix - see Lee et al. Nucl. Acids Res., 6:3073 (1979); Cooney et al., Science, 241:456 (1988); and Dervan et al., Science, 251: 1360 (1991), thereby preventing transcription and the production of GFRQ-4. The antisense RNA oligonucleotide hybridises to the mRNA in vivo and blocks translation of an mRNA molecule into the GFRQ-4 receptor.

Antibodies to the GFR α -4 receptor according to the invention are also provided which may be used in a medicament or in a pharmaceutical composition.

Antibodies to the GFRα-4 of the invention may,
advantageously, be prepared by techniques which are known in the art. For example, polyclonal antibodies may be prepared by inoculating a host animal, such as a mouse, with the polypeptide according to the invention or an epitope thereof and recovering immune
serum. Monoclonal antibodies may be prepared according to known techniques such as described by Kohler R. and Milstein C., Nature (1975) 256, 495-497.

Antibodies according to the invention may also be used in a method of detecting for the presence of GFRα-4 by reacting the antibody with a sample and identifying any protein bound thereto. A kit may also be provided for performing said method which comprises an antibody according to the invention and means for reacting the antibody with said sample.

Advantageously, the antibody according to the invention may also be used as a medicament or in the preparation of a medicament for treating diseases associated with expression of the $\mbox{GFR}\alpha\mbox{-4}$ of the invention. The invention also further provides a pharmaceutical composition comprising said antibody together with a pharmaceutically acceptable carrier, diluent or excipient therefor.

Proteins which interact with the polypeptide of the 10 invention may be identified by investigating proteinprotein interactions using the two-hybrid vector system first proposed by Chien et al (1991), Proc. Natl. Acad. Sci. USA 88 : 9578-9582.

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This technique is based on functional reconstitution in vivo of a transcription factor which activates a reporter gene. More particularly the technique comprises providing an appropriate host cell with a DNA construct comprising a reporter gene under the 20 control of a promoter regulated by a transcription factor having a DNA binding domain and an activating domain, expressing in the host cell a first hybrid DNA sequence encoding a first fusion of a fragment or all of a nucleic acid sequence according to the invention 25 and either said DNA binding domain or said activating domain of the transcription factor, expressing in the host at least one second hybrid DNA sequence, such as, a library or the like, encoding putative binding proteins to be investigated together with the DNA 30 binding or activating domain of the transcription factor which is not incorporated in the first fusion; detecting any binding of the proteins to be investigated with a protein according to the invention by detecting for the presence of any reporter gene 35 product in the host cell; optionally isolating second hybrid DNA sequences encoding the binding protein.

excipient therefor.

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Proteins which bind to the $GFR\alpha-4$ receptor can be identified using this technique. The proteins identified can also be used to identify compounds which acts as agonists/antagonists of these proteins. The structure of the receptor can also be used to design agonists or antagonists of the receptor. The present invention also comprises an agonist or antagonist of the human $GFR\alpha-4$ receptor according to the invention which agonist or antagonist advantageously may also be used as a medicament or in a pharmaceutical composition together with a pharmaceutically acceptable carrier diluent or

- Agonists or antagonists may be identified by 15 contacting a cell expressing $GFR\alpha\text{--}4$ with a compound to be tested and monitoring the degree of any $\mbox{GFR}\alpha\mbox{-}4$ mediated functional or biological response, such as for example, by monitoring the level of
- phosphorylation in said cell or by cytosensor or 20 ligand binding assays in the presence of cRET or similar proteins in the signal transduction pathway. Preferably, the cell may be a host cell or transgenic cell according to the invention as defined herein. Agonists and antagonists of $GFR\alpha-4$ may also be
- 25 identified by, for example, contacting a membrane preparation comprising $GFR\alpha\text{--}4$ with the compound to be tested in the presence of cRET or other similar proteins involved in the signal transduction pathway of which $GFR\alpha-4$ is a component and monitoring the 30 interaction of $GFR\alpha-4$ with cRET or said similar proteins. Advantageously, any compounds or molecules identified as agonists or antagonists in relation to $\mathsf{GFR}\alpha\text{--}4$ may themselves be used in a pharmaceutical composition as defined above or as a medicament.

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Also provided by the invention are molecules or compounds that act on the signal transduction pathway of which GFR α -4 or a functional equivalent belong. Alternatively, the molecules may interfere with complex formation or interaction of GFR α -4 or its functional equivalent, with cRET or a similar protein in the signal transduction pathway of which GFR α -4 is a component.

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- 10 Furthermore, the present invention relates to a method of producing an antagonist or agonist of $GFR\alpha-4$ according to the invention comprising the steps of any one of the above described screening methods; and additionally
 - (i) synthesizing the compound obtained or identified in said method or an physiologically acceptable analog or derivative thereof in an amount sufficient to provide said antagonist or agonist in a therapeutically effective amount to a patient; and/or
 - (ii) combining the compound obtained or identified in said method or an analog or derivative thereof with a pharmaceutically acceptable carrier".

The compounds isolated by the above methods also serve as lead compounds for the development of analog compounds. The analogs should have a stabilized electronic configuration and molecular conformation that allows key functional groups to be presented to the GFR α -4 receptor in substantially the same way as the lead compound. In particular, the analog compounds have spatial electronic properties which are comparable to the binding region, but can be smaller molecules than the lead compound, frequently having a molecular weight below about 2 kD and preferably below

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about 1 kD. Identification of analog compounds can be performed through use of techniques such as selfconsistent field (SCF) analysis, configuration interaction (CI) analysis, and normal mode dynamics analysis. Computer programs for implementing these 5 techniques are available; e.g., Rein, Computer-Assisted Modeling of Receptor-Ligand Interactions (Alan Liss, New York, 1989). Methods for the preparation of chemical derivatives and analogues are well known to those skilled in the art and are 10 described in, for example, Beilstein, Handbook of Organic Chemistry, Springer edition New York Inc., 175 Fifth Avenue, New York, N.Y. 10010 U.S.A. and Organic Synthesis, Wiley, New York, USA. Furthermore, said derivatives and analogues can be tested for their 15 effects according to methods known in the art; see also supra. Furthermore, peptidomimetics and/or computer aided design of appropriate derivatives and analogues can be used.

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Compounds identified as agonists or antagonists in relation to $\text{GFR}\alpha\text{--}4$ or as ligands or compounds which interfere with the signal transduction pathway of which $GFR\alpha-4$ is a part, may advantageously be used in the preparation of a medicament for treatment of 25 neurodegenerative diseases, such as, for example, Alzheimers disease, Parkinsons disease, Motor Neuron Disease, peripheral neuropathy, spinal cord injury, familial hirschsprung disease, in addition to various carcinomas such as for example in gastrointestinal 30 cancer and also in treatment of diseases which may be associated with $GFR\alpha-4$ dysfunction. Compounds identified as antagonists may, advantageously, be used in the preparation of a medicament for the treatment of carcinoma or in alleviating pain. 35

The present invention also further comprises a method of identifying ligands of $GFR\alpha-4$ according to the invention, which method comprises contacting said receptor with either a cell extract or alternatively a compound to be tested for its potential as a $GFR\alpha-4$ ligand, and isolating any molecules bound to $GFR\alpha-4$.

A diagnostic kit is also provided by the present invention, which kit, comprises a probe including any of, a nucleic acid molecule encoding a GFR α -4 protein 10 according to the invention, a molecule capable of hybridising thereto under high stringency conditions, a fragment of said nucleic acids, an antisense molecule according to the invention, together with means for contacting biological material to be tested 15 with said nucleic acid probe. A diagnostic kit in accordance with the invention may also comprise an agonist or antagonist in relation to $GFR\alpha-4$ or an antibody, preferably a monoclonal antibody to $\mathsf{GFR}\alpha\text{--}4$. Thus, advantageously, the kit may be used, as 20 appropriate to identify, for example, cells expressing or lacking in said receptor or genetic defects or the like or for determining whether a compound is a agonist or an antagonist of $GFR\alpha-4$ receptor. Kits for determining whether a compound is an agonist or an 25 antagonist in relation to $\mathsf{GFR}\alpha\text{--}4$ may comprise a cell or membrane preparation expressing said receptor according to the present invention, means for contacting said cell with said compound and means for monitoring the level of any $\mbox{GFR}\alpha\mbox{--}4$ mediated functional 30 or biological response, by for example measuring the level of phosphorylation in said cell or by cytosensor or ligand binding assays in the presence of cRET or similar proteins involved in the signal transduction pathway of which GFR α -4 is a component. 35

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The present invention may be more clearly understood from the following exemplary embodiment with reference to the accompanying figures wherein;

Figure 1: is an illustration of the Structure of the 5 rat $GFR\alpha-4$ gene. The top line shows a scale in bp. The line below shows the genomic structure of the rat $GFR\alpha-4$ gene. Exons are represented by boxes and numbered, intron sequences are depicted as lines. The sizes (in bp) of introns and exons are indicated above 10 the diagram. The translation start codon is indicated by an arrow and the stop codon by an asterisk. The cDNA sequences of variants A and B obtained by splicing out the intron sequences is shown below the genomic sequence. Alternative splicing of intron 5 15 results in an earlier stop codon in splice variant B. The predicted protein sequences of variants A and B are shown at the bottom. The predicted signal peptide, a putative N-glycosylation site and a hydrophobic COOH-terminal region preceded by one or two possible 20 sites for GPI-cleavage (in variant A only) are indicated on the diagrams.

Figure 2: is an alignment of the predicted protein sequences of splice variants A and B of rat $GFR\alpha-4$. The sequences of rat $GFR\alpha-4$ splice variants A and B were aligned using the ClustalW alignment program (EMBL, Heidelberg, Germany). Amino acid residues conserved between the 2 variants are included in the black areas. Amino acid residues are numbered to the right. The dashes indicate gaps introduced into the sequence to optimize the alignment.

Figure 3: is an alignment of the predicted protein sequences of GFR α family members. The sequence of rat GFR α -4 variants A and B, rat GFR α -1 (EMBL acc. no. U59486), rat GFR α -2 (EMBL acc. no. AF003825), mouse

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GFR α -3 (EMBL acc. no. AB008833) and chicken GFR α -4 (EMBL acc. no. AF045162) were aligned using the ClustalW alignment program (EMBL, Heidelberg, Germany). Amino acid residues conserved between all 6 proteins are included in the black areas. Residues conserved between 4 or 5 of the sequences are shaded in grey. Cysteine residues conserved between all six GFR α 's are indicated with an asterisk above the sequence. Amino acid residues are numbered to the right. The dashes indicate gaps introduced into the sequence to optimize the alignment.

Figure 4: Northern blot analysis of rodent GFR α -4 mRNA expression. The expression of rat and mouse GFR α -4 mRNA in different tissues was assessed using a probe corresponding to the coding sequence of rat GFR α -4 to analyze blots of poly(A)-rich RNA. (A) Rat Multiple Tissue Northern (MTN) blot; (B) Mouse MTN blot and (C) Mouse Embryo MTN blot. Apparent sizes are indicated (in kilobase pairs) by horizontal lines to the left of each panel.

Figure 5. The rat GFR α -4 gene is localized on chromosome 3q36. A mixture of two rat GFR α -4 probes was used for FISH analysis. (A) Double-spot FISH signals on the middle-distal part of rat chromosome 3 (arrows). (B) Position of the GFR α -4 gene locus on rat chromosome 3q36.

30 Oligonucleotide synthesis for PCR and DNA sequencing.

All oligonucleotide primers used were ordered from Eurogentec (Seraing, Belgium). Insert-specific sequencing primers (15- and 16-mers) and primers for use in PCR reactions were designed manually. DNA was prepared on Qiagen-tip-20 or -100 anion exchange or Qiaquick spin columns (Qiagen GmbH, Düsseldorf,

Germany) and recovered from the columns in 30 µl TE-buffer (10 mM Tris.HCl, 1 mM EDTA (sodium salt), pH 8.0). Sequencing reactions were done on both strands using the ABI prism BigDye Terminator Cycle sequencing kit and were run on an Applied Biosystems 377XL sequencer (Perkin Elmer, ABI Division, Foster City, CA, USA). The Sequencher™ software was used for sequence assembly and manual editing (GeneCodes, AnnArbor, MI, USA).

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Identification of a cDNA sequence encoding a novel member of the GFR family

Using the human GFR α -1, GFR α -2 or GFR α -3 DNA or protein sequences as the query sequence, BLAST (Basic Local Alignment Search Tool; Altschul et al., 1990) searches were performed on the daily updates of the public EMBL database. A mouse EST (expressed sequence tag) sequence with EMBL accession number AU035938 showed homology to GFR α -1, GFR α -2 and GFR α -3. The smallest sum probabilities (SSP) obtained by the BLAST analyses are summarized in Table 1.

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Table 1: BLAST	results.	
Query sequence	DNA/PROTEIN	SSP
GFRα-1	protein	7.5e-25
GFRα-2	protein	1.3e-12
GFRα-3	protein	2.2e-20
GFRα-1	DNA	6. 6 e-09
GFRα-2	DNA	>0.011
GFRα-3	DNA	0.0096

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AU035938 (sequence 1) is a sequence of 792 bp derived from a mouse brain cDNA library. To obtain consistent homology with other members of the GFR α family upon translation a frame shift has to be introduced near position 165 in the DNA sequence. It is not clear whether this is due to a sequencing error or whether there is another explanation. Using this EST sequence

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as the query sequence, the BLAST search against the public EMBL database was repeated. One additional clone (acc. no. AA823200; sequence 2) yielded a significant SSP of le-18. Upon inspection of this 497 bp clone, which was derived from a mouse mammary gland cDNA library, only the first 61 bp were identical with part of AU035938 (position 353 to 415). The rest of the sequence of AA823200 was different from AU035938, but contained parts of which the translated amino acid sequence showed homology with the other GFRa's. Therefore it was hypothesized that AU035938 and AA823200 could represent two variant forms of the same receptor, which was called GFRa-4.

15 Cloning of mouse GFRα-4 cDNA

First, we tried to amplify a fragment of the mouse $GFR\alpha-4$ cDNA on Marathon ReadyTM cDNAs (Clontech Laboratories, Palo Alto, CA, USA) derived from mouse brain and mouse embryo. Primers were designed using the EST sequences (EMBL acc. no. AU035938 and AA823200) to amplify a 274 bp fragment of mouse $GFR\alpha-4$. The primers used for the amplification of mouse $GFR\alpha-4$ are shown in the table below.

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Table 2: Primers used for the amplification of mouse $GFR\alpha-4$ DNA sequences.

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Name	Sequence	n
MOUSE-GFRa4-sp2	CGCGTTGTCTGCGCGTCTACG	21
MOUSE-GFRa4-sp3	CGGCGCGAAGAATGCGAAGC	20
	CACCCACGTACCATGGCATGTGC	23
MOUSE-GFRα4-ap2	CACCCACOTAGGATTE	

PCR reactions were done using the Taq polymerase system (Boehringer Mannheim, Mannheim, Germany). PCR reactions were performed in a total volume of 50 µl, containing 1x Taq PCR reaction buffer, 0.25 mM dNTP,

0.5 μM of primers MOUSE-GFR $\alpha 4$ -sp2 and MOUSE-GFR $\alpha 4$ -ap2, 1 µl of Tag polymerase and 2 µl of mouse embryo or mouse brain Marathon Ready $^{\text{TM}}$ cDNA. Samples were heated to 95°C for 5 min and cycling was done for 30 s at 94°C, 1 min at 60°C and 45 s at 72°C for 35 cycles, 5 with a final step of 7 min at 72°C. A semi-nested PCR was then performed on 1 µl of the primary PCR reaction with primers MOUSE-GFR $\alpha4$ -sp3 and MOUSE-GFR $\alpha4$ -ap2. PCR reactions were performed in a total volume of 50 μ l, containing 1x Taq PCR reaction buffer, 0.25 mM dNTP, 10 0.5 μM of primers MOUSE-GFR $\alpha 4$ -sp3 and MOUSE-GFR $\alpha 4$ -ap2, 1 μl of Taq polymerase and 1 μl of primary PCR product. Samples were heated to 95°C for 5 min and cycling was done for 30 s at 94°C, 1 min at 60°C and 45 s at 72°C for 35 cycles, with a final step of 7 min15 at 72°C. PCR products were analysed on a 1% (w/v)agarose gel in 1x TAE buffer (40 mM Tris-acetate, 1 mM EDTA (sodium salt), pH 8.3). A PCR fragment of the expected size (270 bp) was excised from the gel and purified with the Qiaquick gel extraction kit (Qiagen 20 Gmbh, Hilden, Germany). The PCR fragments were sequenced to confirm their identity. The obtained sequence corresponded to the EST database sequences. In order to determine the upstream and downstream coding sequences of mouse GFR α -4, 5' and 3' RACE 25 experiments were performed. Since these experiments did not work as expected and since, at some points, frame shifts had to be introduced in the mouse $\mathsf{GFR}\alpha\text{--}4$ sequence to yield consistent homology with other $\mbox{\rm GFR}\alpha's$ after translation, we decided to shift to the 30 cloning of the rat homologue of mouse $GFR\alpha-4$.

Identification and cloning of rat GFRα-4 cDNA sequences

The cDNA sequences with accession number AU035938 and AA823200 described above were used as the query sequence in BLAST searches on the proprietary LifeSeq and ZooSeq databases (Incyte Pharmaceuticals, Palo 5 Alto, CA, USA). Two rat clones with high homology to the mouse $GFR\alpha-4$ sequences were identified: number 701290919H1 (270 bp; hit with AU035938 (SSP = 1.1e-32) and with AA823200 (SSP = 1.3e-21)) and number 701291473H1 (250 bp; hit only with AA823200 (SSP = 10 4.3e-42)). From comparing the translated protein sequences derived from clones 701291473H1 and 701290919H1 to the known GFRa protein sequences, it could be deduced that sequence 701290919H1 was probably localised 5' to sequence 701291473H1 and that 15 these sequences were almost adjacent to each other in the full GFR α -4 cDNA sequence. Therefore, two forward primers (RAT-GFR α 4-sp1 and RAT-GFR α 4-sp2) were designed in the 5' region of sequence 701290919H1 and two reverse primers (RAT-GFR α 4-apl and RAT-GFR α 4-ap2) 20 in the 3' region of sequence 701291473H1. All primer sequences used in PCR experiments are summarized in Table 3.

Table 3: Primers used for the amplification of rat $GFR\alpha-4$ sequences. The RACE-apl and RACE-ap2 primers are included in the Marathon ReadyTM cDNA kit.

Name	Sequence	n
RAT-GFRα4-sp1	GTGGTCACCCCCAACTACCTGG	22
RAT-GFRα4-sp2	GCCTTCCGCAAGCTTTTTACAAGG	24
RAT-GFRa4-sp3	GCTCTTCTGCGGATGCGAAGGC	22
RAT-GFRα4-sp4	AGCTGCCGGGTTTACTGATGCTAC	24
RAT-GFRa4-sp5	GATGCTACTCTCCCAAGGTCAGGC	24
RAT-GFRa4-sp6	CTGGTAAGCTTTAAGGCAGAGGAGACC	27
RAT-GFRα4-ap1	CATGGCAGTCAGCTGTGTTGTCC	23
RAT-GFRα4-ap2	CAGCTGTGTTGTCCATGGTTCACC	24
RAT-GFRα4-ap3	TGGTTGCGAGCTGTCAAAGGCTTGTATGGC	30
RAT-GFRα4-ap4	GGGGTTCCTTGTAAAAAGCTTGCGGAAGGC	30
RAT-GFRα4-ap5	GGTCCAAGGCTTCAGGCAGGAAGG	25
RAT-GFRα4-ap6	GCCTTCGCATCCGCAGAAGAGC	22
RAT-GFRα4-ap7	CCAGGTAGTTGGGGGTGACCACG	23
RAT-GFR04-ap7b	CCCAGGCATTGCGCCACGTA	20
RAT-GFRα4-ap8	CATTGCGCCACGTACTCGGAGC	22
RAT-GFRα4-ap9	GACCTGAGGGCAAGGGAGTTTCA	23
RAT-GFRα4-ap10	GCAAGGGAGTTTCAGTTCAGTGAGC	25
RACE-ap1	CCATCCTAATACGACTCACTATAGGGC	2
RACE-ap1	ACTCACTATAGGGCTCGAGCGGC	2:

A PCR was then performed using primers RAT-GFRα4-spl and RAT-GFRα4-apl on rat brain Quickclone cDNA (Clontech Laboratories, Palo Alto, CA, USA) to confirm the presence of rat GFRα-4 in brain-derived cDNA. Since the DNA sequence coding for the rat GFRα-4 sequence has a high G+C content in this region, PCR reactions were done using the Advantage-GC PCR kit (Clontech). PCR reactions were performed in a total volume of 50 μl, containing 1x GC cDNA PCR reaction buffer, 0.2 mM dNTP, 1 M GC-MELTä, 200 nM of primers RAT-GFRα4-spl and RAT-GFRα4-apl, 1 μl of Advantage KlenTaq polymerase mix and 1 μl of rat brain Quickclone cDNA. Samples were heated to 95°C for 1 min and cycling was done for 1 min at 95°C, 1 min at 56°C

and 1 min at 72°C for 30 cycles, with a final step of 7 min at 72°C. A nested PCR was then performed on 1 μl of the primary PCR reaction with primers RAT-GFR $\alpha4$ -sp2 and RAT-GFR α 4-ap2. PCR reactions were performed in a total volume of 50 µl, containing 1x GC cDNA PCR 5 reaction buffer, 0.2 mM dNTP, 1 M GC-MELTä, 200 nM of primers RAT-GFR α 4-sp2 and RAT-GFR α 4-ap2, 1 μ l of Advantage KlenTag polymerase mix and 1 µl of primary PCR product. Samples were heated to 95°C for 1 min and cycling was done for 30 s at 95°C, 1 min at 56° C and 1 10 min at 72° C for 25 cycles, with a final step of 7 min at 72°C. PCR products were analysed on a 1% (w/v) agarose gel in 1x TAE buffer (40 mM Tris-acetate, 1 mM EDTA (sodium salt), pH 8.3). Two PCR fragments of approximately 1100 and 200 bp, respectively, were 15 excised from the gel and purified with the Qiaquick gel extraction kit (Qiagen Gmbh, Hilden, Germany). The PCR fragments were sequenced to confirm their identity. The smallest fragment yielded a sequence of 211 bp corresponding to the joined sequences 20 701290919Hl and 701291473Hl. The larger fragment yielded a sequence of 1049 bp of which 18 bp at the 5' end, 59 bp at the 3^{\prime} end and an internal stretch of 92 bp corresponded to the sequence of the 211 bp fragment, but which had additional sequence stretches 25 in between. This fragment represented a variant of rat $GFR\alpha-4$.

Both clones 701291919H1 and 701291473H1 were obtained from Incyte Pharmaceuticals and the inserts completely sequenced. The sequences are included in this application (sequence 3 = 701290919H1 and sequence 4 = 701291473H1). Both clones were derived from the same 7-day old rat brain cortex cDNA library. Both clones

differ in their 5' ends (first 134 bp in 701291473H1 and first 227 bp in 701290919H1) but are identical thereafter. Both contain part of the $GFR\alpha-4$ coding sequence up to a stop codon (position 184-186 in 701291473H1 and 277-279 in 701290919H1). A 3' 5 untranslated region of 549 bp followed by a poly(A)tail is then present in both clones. We hypothesized that both clones are different variants of the rat GFR α -4 gene. Primers (RAT-GFR α 4-ap3 and RAT-GFR α 4-ap4) were designed on a part of the sequence common to both 10 variants to perform 5' RACE experiments in order to determine the 5' end of the rat $GFR\alpha-4$ cDNA. First, a 5' RACE PCR was performed on rat brain Marathon Ready $^{\text{TM}}$ cDNA (Clontech). PCR reactions were performed in a total volume of 50 µl, containing 1x GC 15 cDNA PCR reaction buffer, 0.2 mM dNTP, 1 M or 1.5 M GC-MELTä, 200 nM of primers RAT-GFRα4-ap3 and RACEapl, 1 μl of Advantage KlenTaq polymerase mix and 5 μl of rat brain Marathon Ready $^{\text{TM}}$ cDNA. Samples were heated to $95\,^{\circ}\text{C}$ for 30 s and cycling was done for 30 s 20 at 95°C, 4 min at 72°C for 5 cycles, then 30 s at 95°C, 4 min at 70°C for 5 cycles, then 30 s at 95°C, 4 min at 68° C for 25 cycles, with a final step of 7 min at 68°C. A nested PCR was then performed on 1 μl of the primary PCR reaction with primers RAT-GFR $\alpha4$ -ap4 25 and RACE-ap2. PCR reactions were performed in a total volume of 50 µl, containing 1x GC cDNA PCR reaction buffer, 0.2 mM dNTP, 1 M or 1.5 M GC-MELTä, 200 nM of primers RAT-GFR α 4-ap4 and RACE-ap2, 1 μ l of Advantage KlenTaq polymerase mix and 1 μl of primary PCR 30 product. Cycling was done using exactly the same parameters as for the primary PCR. PCR products were analysed on a 1% (w/v) agarose gel in 1x TAE buffer (40 mM Tris-acetate, 1 mM EDTA (sodium salt), pH 8.3).

A fragment of approximately 350 bp was excised from the gel and cloned into the plasmid vector pCR2.1-TOPO using the TOPO TA cloning kit according to manufacturer's instructions (Invitrogen BV, Leek, The Netherlands). One of the resulting clones yielded an 5 insert sequence of 387 bp which extended the rat $GFR\alpha$ -4 sequence in the 5' direction. Upon translation, this additional cDNA sequence yielded a protein sequence without any internal stop codons and with substantial homology to the other known GFR α sequences. Since no 10 putative ATG start codon could be detected within this additional sequence, novel primers (RAT-GFR α 4-ap5 and RAT-GFR α 4-ap6) were designed at the 5' end of this sequence to perform additional 5' RACE experiments. First, a 5' RACE PCR was performed on rat heart 15 Marathon Ready $^{\text{TM}}$ cDNA (Clontech). PCR reactions were performed in a total volume of 50 μ l, containing 1x GC cDNA PCR reaction buffer, 0.2 mM dNTP, 1 M GC-MELTä, 200 nM of primers RAT-GFR $\alpha 4$ -ap5 and RACE-ap1, 1 μl of Advantage KlenTaq polymerase mix and 5 µl of rat heart 20 Marathon Ready $^{\text{TM}}$ cDNA. Samples were heated to 95 $^{\circ}$ C for 30 s and cycling was done for 30 s at 95°C, 4 min at 72°C for 5 cycles, then 30 s at 95°C , 4 min at 70°C for 5 cycles, then 30 s at 95°C, 4 min at 68°C for 25 $\,$ cycles, with a final step of 7 min at 68°C. A nested 25 PCR was then performed on 1 μl of the primary PCR reaction with primers RAT-GFR α 4-ap6 and RACE-ap2. PCR reactions were performed in a total volume of 50 μ l, containing 1x GC cDNA PCR reaction buffer, 0.2 mM dNTP, 1 M GC-MELTä, 200 nM of primers RAT-GFRα4-ap6 30 and RACE-ap2, 1 µl of Advantage KlenTaq polymerase mix and 1 μ l of primary PCR product. Cycling was done using exactly the same parameters as for the primary PCR. PCR products were analysed on a 1% agarose gel. A

fragment of approximately 200 bp was excised from the gel and cloned into the plasmid vector pCR2.1-TOPO using the TOPO TA cloning kit as described above. Sequencing of two resulting clones extended the rat $GFR\alpha-4$ sequence with another 128 bp in the 5' 5 direction. Based on this sequence, another primer set (RAT-GFR α 4-ap7 and RAT-GFR α 4-ap8) was designed to perform additional 5' RACE experiments. RACE PCR was performed on rat brain, heart and kidney Marathon $Ready^{TM}$ cDNA. PCR reactions were performed in a total 10 volume of 50 µl, containing 1x GC cDNA PCR reaction buffer, 0.2 mM dNTP, 1 M GC-MELTä, 200 nM of primers RAT-GFR α 4-ap7b and RACE-ap1, 1 μ l of Advantage KlenTaq polymerase mix and 5 μl of rat heart, brain or kidney Marathon Ready $^{\text{TM}}$ cDNA. Samples were heated to 95°C for 15 30 s and cycling was done for 30 s at 95°C, 4 min at 72°C for 5 cycles, then 30 s at 95°C, 4 min at 70°C for 5 cycles, then 30 s at 95°C, 4 min at 68°C for 25 cycles, with a final step of 7 min at 68°C. A nested PCR was then performed on 1 μl of the primary PCR 20 reaction with primers RAT-GFRlpha4-ap8 and RACE-ap2. PCR reactions were performed in a total volume of 50 μ l, containing 1x GC cDNA PCR reaction buffer, 0.2 mM dNTP, 1 M GC-MELTä, 200 nM of primers RAT-GFR α 4-ap8 and RACE-ap2, 1 µl of Advantage KlenTaq polymerase mix 25 and 1 μ l of primary PCR product. Cycling was done using exactly the same parameters as for the primary PCR. PCR products were analysed on a 1% agarose gel. Several fragments ranging in size from approximately 200 bp to 1200 bp were visible on the gel and were 30 excised and cloned in vector pCR2.1-TOPO using TOPO-TA cloning. The inserts of several clones were sequenced. From these clones, the sequence of rat GFRlpha-4 could be extended in the 5' direction. Two different sequences

were identified. One sequence extended the ratGFR α -4 sequence with 215 bp in the 5' direction and included an in-frame start codon preceded by an in-frame upstream stop codon. The resulting predicted protein sequence (52 additional amino acid residues) includes 5 a predicted signal peptide of 29 amino acid residues (as determined by the SPScan program included in the Wisconsin package version 10.0, Genetics Computer Group (GCG), Madison, Wisconsin, USA; score 7.0, probability 1.171e-02). The other sequence determined 10 by these 5' RACE experiments extended the ratGFR α -4 sequence with 552 bp in the 5' direction and also included an in-frame start codon preceded by an inframe upstream stop codon. The most 3' 79 base pairs of this novel sequence were identical to the 3' 79 15 base pairs of the 215 bp sequence, but the rest of the sequence was different. The resulting predicted protein sequence (113 additional amino acid residues), however, did not have a predicted signal peptide sequence at the $\mathrm{NH}_2\text{-terminus}$ (SPScan, GCG package). 20 The different partial cDNA sequences resulting from the subsequent 5' RACE experiments together with the sequences from the Incyte database were compared and merged into several possible rat $\mbox{GFR}\alpha\mbox{-4}$ variants. In order to identify which of the identified variants are 25 real, primers were designed 5' of the translation start codon (primers RAT-GFR $\alpha4$ -sp4 and RAT-GFR $\alpha4$ -sp5 for the "long" 5' variant resulting from the 552 bp RACE fragment and RAT-GFRα4-sp6 for the "short" 5' variant resulting from the 215 bp RACE fragment) and 30 3^{\prime} of the translation stop codon (RAT-GFR $\alpha4$ -ap9 and RAT-GFRlpha4-ap10). These primers were then used to amplify the full GFR α -4 coding sequences using cDNA derived from different rat tissues.

First, sequences coding for the "long" 5' variant were amplified by PCR. PCR reactions were performed in a total volume of 50 μ l, containing 1x GC cDNA PCR reaction buffer, 0.2 mM dNTP, 1 M GC-MELTä, 200 nM of primers RAT-GFR α 4-sp4 and RAT-GFR α 4-ap9, 1 μ 1 of 5 Advantage KlenTaq polymerase mix and 5 µl of rat heart, brain or kidney Marathon Ready TM cDNA. Samples were heated to 95°C for 1 min and cycling was done for $45 \text{ s at } 95^{\circ}\text{C}$, 1 min at 57°C and 1 min at 72°C for 35cycles, with a final step of 7 min at 72°C. A nested 10 PCR was then performed on the primary PCR reaction with primers RAT-GFR $\alpha4$ -sp5 and RAT-GFR $\alpha4$ -ap10. PCR reactions were performed in a total volume of 50 μ l, containing 1x GC cDNA PCR reaction buffer, 0.2 mM dNTP, 1 M GC-MELTä, 200 nM of primers RAT-GFRα4-sp5 15 and RAT-GFR α 4-ap10, 1 μ 1 of Advantage KlenTaq polymerase mix and 1 µl of primary PCR product. Cycling was done using exactly the same parameters as for the primary PCR, except that 30 PCR cycles were done instead of 35. PCR products were analysed on a 1% 20 agarose gel. Several fragments ranging in size from approximately 1000 to 1250 bp were excised from gel and cloned in vector pCR2.1-TOPO using TOPO-TA cloning. The inserts of several clones were sequenced. Next, sequences coding for the "short" 5' variant were 25 amplified by PCR. PCR reactions were performed in a total volume of 50 µl, containing 1x GC cDNA PCR reaction buffer, 0.2 mM dNTP, 1 M GC-MELTä, 200 nM of primers RAT-GFR $\alpha 4$ -sp6 and RAT-GFR $\alpha 4$ -ap9, 1 μl of Advantage KlenTaq polymerase mix and 5 µl of rat heart 30 Marathon Ready $^{\text{TM}}$ cDNA. Samples were heated to 95°C for 5 min and cycling was done for 30 s at 94°C, 1 min at 57°C and 2 min 30 s at 72°C for 35 cycles, with a final step of 7 min at 72°C. PCR products were

analysed on a 1% agarose gel. Several fragments ranging in size from approximately 1500 to 2200 bp were excised from gel and cloned in vector pCR2.1-TOPO using TOPO-TA cloning. The inserts of several clones were sequenced. Analysis of all the obtained sequences (16 resulting clones were completely sequenced) allowed the rat $GFR\alpha-4$ DNA sequence to be divided into 6 sequence stretches common to all identified variants, with 5 intervening sequence stretches present or absent depending on the variant. All 5 10 intervening sequences contain 5' and 3' splice site consensus sites (GT at the 5' end and AG at the 3' end of the intron sequence) (Senapathy et al., 1990) (see table 4 below) and could thus potentially represent unspliced introns. 15

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In order to strengthen the hypothesis that the identified variants could result from the conservation of unspliced introns in certain mRNA transcripts, the rat $GFR\alpha-4$ sequence was compared to the genomic 20 sequence of human GFR α -1 (Angrist et al., 1998). From this analysis, it was apparent that the $GFR\alpha-4$ sequences common to all transcripts coincided with exons in $GFR\alpha-1$ (see table 4 below). The intervening sequences absent in some transcripts coincided with 25 intron sequences in human GFR α -1. Therefore, we considered all intervening sequences as unspliced introns. The intron present between exon 5 and exon 6 can be spliced out in two different ways and results in the presence of two different splice variants of 30 rat $GFR\alpha-4$, which we have called variant A and variant в.

Sequence 5 shows the consensus sequence for rat $\mbox{GFR}\alpha\mbox{-}4$ including the intron sequences (intron 1: bp 125 to 684; intron 2: bp 1040 to 1088; intron 3: bp 1199 to 1278; intron 4: bp 1414 to 2154; intron 5A: bp 2247 to 2385 and intron 5B: bp 2231 to 2314). A polymorphism 5 was detected at position 2244 in sequence 5, with T found in 50% of the sequenced clones and C in the other 50%. This polymorphism leads to an amino acid change in the protein (variant A) from W to R, in the hydrophobic region involved in GPI-anchoring. 10 Figure 1 schematically shows the structure of the rat $\mbox{GFR}\alpha\mbox{-4}$ gene together with the derived cDNA for splice variants A and B after splicing out of the intron sequences and the translated protein sequences of variants A and B with their characteristics. 15 Table 4 shows the DNA sequence at the intron-exon boundaries together with the sizes of identified introns and exons. The right column shows the sizes of the corresponding exons in the genomic sequence of human GFR α -1 (from Angrist et al., 1998). 20

Table 4: Intron-exon structure of rat $GFR\alpha-4$.

					Splice donor	Corresponding
	Exon	Size	Intron	Splice	Sprice donor	GFRα-1 exon
		(bp)	size (bp)	acceptor		· ·
	l	(-,-		1		size (bp)
					GAGgtaaggaggt	
25	1	>124	560	ccctcaccagGGT	CCGqtgcgtgcgg	337
	2	355	49	ecettactagoor	TAGgtacgctggg	110
	3	110	80	gcgcgcgcagGCC		135
	1	135	741	gtccctgcagGCA	TGGgtgagggggc	182
	5	92	139 (varA)	cactccatagATG	CGGgtaggtatgg	1 102
	١	12	84 (varB)		TGGgtgctgtttc	753
30	6	>137		ttgtcccaagGTG	-	/53
50	ľ	1	1	cccttctcagGCA		

The consensus sequence obtained by removing introns 1 to 4 and intron 5A (sequence 6; variant A) translates into a protein of 273 amino acid residues with a

calculated molecular mass of 29.7 kDa and an isoelectric point of 8.92 (sequence 8). The consensus sequence obtained by removing introns 1 to 4 and intron 5B (sequence 7; variant B) translates into a protein of 258 amino acid residues with a calculated 5 molecular mass of 28.0 kDa and an isoelectric point of 8.91 (sequence 9). Figure 2 shows the alignment of variants A and B of rat $GFR\alpha-4$. The protein sequences are both similar to the known GFR α sequences and only differ from each other in a small amino acid stretch 10 at the carboxy-terminus. These two sequences probably represent biologically active $GFR\alpha-4$ variants. Since all the other variants sequenced contain one or more intron sequences, they are probably intermediates of RNA processing. It is not clear why all these 15 intermediates are present in cDNA derived from purified mRNA and why it is so difficult to amplify a cDNA sequence derived from a completely spliced mRNA transcript. GFR α -1 to -4 are characterized by a COOHterminal sequence typical of a glycosyl-phosphatidyl 20 inositol (GPI)-anchored protein, consisting of a hydrophobic region of 17-31 amino acid residues preceded by a hydrophilic sequence containing a stretch of three small amino acids such as Asp, Cys, Ala, Ser, Gly or Asn (Gerber et al., 1992). The rat 25 $GFR\alpha$ -4 variant A protein sequence has a hydrophobic carboxy-terminus of 21 amino acid residues (position 253 to 273) preceded by two possible GPI cleavage sites (DSS at position 234 to 236 or NAG at position 250-252). Variant B has a shorter hydrophilic carboxy-30 terminus, implying that no GPI-anchoring is possible for this variant. This could mean that variant B is a soluble form of the rat $GFR\alpha-4$ receptor. A predicted signal peptide of 29 amino acids is present in both

variants (as determined by the SPScan program included in the GCG package; score 7.0, probability 1.171e-02). In addition, one possible site for N-linked glycosylation (NVS at position 192 to 194 in the protein) is present.

Recently, a model has been proposed for the domain structure of $\mbox{\rm GFR}\alpha's$ based on the comparison of the sequences of mouse GFR α -1 to -3 and chicken GFR α -4 (Airaksinen et al., 1999). The model includes three 10 conserved cysteine-rich domains joined together by less conserved adaptor sequences. The molecules are anchored to the membrane by a GPI-anchor. Rat $\mbox{GFR}\alpha\mbox{-}4$ conforms partly to this model, since it also contains the second and third cysteine-rich region and a 15 possible GPI-anchor (at least for variant A). However, it differs significantly from the other $\mathsf{GFR}\alpha's$ in that the first cysteine-rich region is absent. Figure 3 shows the alignment of rat $GFR\alpha-4$ variants A and B with rat $GFR\alpha-1$ (EMBL acc. no. U59486), rat 20 GFR α -2 (EMBL acc. no. AF003825), mouse GFR α -3 (EMBL acc. no. AB008833) and chicken GFR α -4 (EMBL acc. no. AF045162). The alignment was done using the ClustalW alignment program (EMBL, Heidelberg, Germany). The percentage identity and percentage similarity between 25 members of the GFR α family were calculated by pairwise comparison of the sequences using the GeneDoc software tool (version 2.5.000) and the results are presented in Table 5 below.

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Table 5: % identity and % similarity (between brackets) between members of the GFR α family. Accession numbers of the sequences used in the analysis are mentioned in the text.

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	rGFRa-1	rGFRa-2	mGFRa-3	cGFRa-4	rGFRa-4 (A)	rGFRa-4 (B)
	100	43 (60)	15 (23)	38 (55)	20 (29)	20 (28)
rGFRa-1	100		18 (28)	40 (56)	21 (32)	21 (31)
rGFRa-2		100		16 (25)	22 (30)	20 (29)
mGFRα-3			100		27 (37)	26 (35)
10 cGFRa-4				100		92 (92)
rGFRa-4(A)					100	
-CFD=-4/P)						100

Four members of the GDNF family of neurotrophic factors have been identified so far (GDNF, NTN, PSP, EVN/ARTN). All four signal through binding to a specific GPI-linked GFR α receptor (GFR α -1 for GDNF, GFR α -2 for NTN, GFR α -3 for EVN/ARTN and (chicken) GFR α -4 for PSP) in combination with a common transmembrane tyrosine kinase, cRET. GFR α -4, the coreceptor for PSP, has been identified in chicken only and no mammalian counterpart has been found yet.

The similarity between the rat GFR α -4 described in the present application and the chicken GFR α -4 is 37% (27% identity) suggesting that rat GFR α -4 is a novel member of the GFR α family. GFR α -4 could be the mammalian persephin receptor or, alternatively, could be the receptor for an unidentified GDNF family member.

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Specific binding of persephin to GFRa-4.

Constructs for the expression of soluble GFR α -IgG-Fc fusion proteins were made as follows. cDNA regions of human GFR α -1, GFR α -2 and GFR α -3, chicken GFR α -4 and rat GFR α -4 variant A (coding for amino acid residues 27 to 427, 20 to 431, 28 to 371, 20 to 399 and 29 to

252, respectively), excluding the sequences coding for the signal peptide and for the COOH-terminal hydrophobic region involved in GPI-anchoring, were cloned in-frame in the expression vector Signal pIg plus (R&D Systems Europe Ltd, Abingdon, UK). The 5 inserts of all constructs were confirmed by complete DNA sequence analysis. The resulting proteins expressed from these constructs contain a 17 amino acid residue $\mathrm{NH_2-terminal}$ CD33 signal peptide, the respective $GFR\alpha$ protein region and a 243 amino acid 10 residue COOH-terminal human IgG_1 -Fc fusion domain. Fusion proteins were expressed in CHO cells and purified as described. Chinese hamster ovary (CHO) cells were routinely cultured in DMEM/F12 medium supplemented with 10 % heat inactivated fetal calf 15 serum. Cells were transfected with GFR α -IqGFc fusion constructs using an optimized Lipofectamine Plus method. For this, a total amount of 6.5 µg DNA was incubated with 17.5 μl PLUS reagent in 750 μl serum free medium for 15 min at room temperature. 20 Lipofectamine was diluted 50-fold into serum free culture medium, 750 µl of this mixture was added to the DNA solution. Following a 15 min incubation at room temperature, 3.5 ml serum free medium was added, and the mixture was brought onto the cells (in a 100 25 mm petridish). The cells were incubated for 3h at 37°C in 5 % $\rm CO_2$, after which 5 ml of culture medium, containing 20 % heat inactivated fetal calf serum, was added. 24 h later, the medium was changed into regular culture medium. Transfection efficiencies using these 30 optimized conditions were typically 50-60%. For permanent transfections the selection medium contained either 800 μg G418 or 800 μg G418 and 800 μg hygromycin. Antibiotic resistant clones were expanded

and assayed for expression using specific antibodies. $\mathsf{GFR}\alpha\text{-}\mathsf{IgGFc}$ fusion proteins were purified from the medium of permanently or transiently transfected CHO cells by protein A chromatography. Bound protein was eluted with 0.1 M Na-citrate, pH 3.0 and collected 5 into 1 M Tris buffer, pH 8.4 (dilution ratio 1:6). Protein concentration was estimated by absorbance at 280 nm using an extinction coefficient of 1.5. Surface plasmon resonance (SPR) experiments were performed at 25°C using a BIACORE 3000 instrument (Biacore AB, 10 Uppsala, Sweden). Sensor chip CM5, the amine coupling kit and buffers used were also obtained from Biacore AB. Recombinant PSP, NTN, EVN/ART and GDNF were used as immobilised ligands. Recombinant human GDNF was obtained from R&D Systems Europe Ltd. (Abingdon, UK). 15 $\mathrm{NH_{2}\text{-}terminally}$ 6His-tagged recombinant human NTN, rat PSP and human EVN/ART were produced in E. coli as described previously (Creedon et al., 1997). The carboxylated matrix of a CM5 sensor chip was first activated with a 1:1 mixture of 400 mM N-ethyl-N'-(3-20 dimethylaminopropyl)-carbodiimide and 100 mM Nhydroxy-succinimide for 10 min. Recombinant neurotrophic factors were applied onto the activated surface in 10 mM sodium acetate buffer, pH 4.5 at a flow rate of 5 μ l/min. Unreacted carboxyl groups were 25 blocked with 1 M ethanolamine-HCl. For binding experiments, soluble $GFR\alpha$ -IgGFc fusion proteins were superfused using the kinject program at 30 µl/min. Concentrations of GFR α -IgGFc used in kinetic experiments were between 1 and 100 nM in Hepes 30 buffered saline (150 mM NaCl, 3.5 mM EDTA sodium salt, 0.005% polysorbate 20, 10 mM Hepes, pH 7.4). The association of the $\ensuremath{\mathsf{GFR}\alpha}$ receptors to the immobilised ligands was monitored for 3 min and the dissociation

for 1 min, followed by regeneration with 10 mM glycine buffer. Dissociation was initiated by superfusion with Hepes buffered saline. To improve the quality of sensor data, double referencing was used (Myszka, 1999). Data were analyzed using a global analysis with 5 the BIACORE evaluation software (version 3.0.1). . Global analysis calculates the association rate $(k_{\rm a})$ and dissociation rate $\left(k_{d}\right)$ simultaneously and the apparent equilibrium dissociation constant (K_n) is then calculated as $k_{\rm d}/k_{\rm a}.$ A simple 1:1 Langmuir model 10 was used to fit the data. Specific binding to PSP could be detected with both rat and chicken $\mathsf{GFR}\alpha\text{-}4\text{-}$ IgGFc fusion proteins. The observed binding of $GFR\alpha 4-$ IgGFc was specific as there was no binding to GDNF, NTN or EVN/ART. Control experiments confirmed binding 15 of GFR-1 to GDNF, of GFR α -2 to NTN and of GFR α -3 to EVN/ART. From the binding curves obtained using three determinations at differing concentrations of rat and chicken GFR $\alpha4$ -IgGFc, the binding constants $k_{\rm a}$ (association rate) and k_{d} (dissociation rate) were 20 derived (Table 6).

Table 6: Persephin binding to chicken GFR α -4 and rat GFR α -4.

Binding constants for chicken $GFR\alpha-4-IgGFc$ and rat $GFR\alpha-4-IgGFc$ binding to immobilised persephin as determined by SPR. The mean association rate (k_a) , dissociation rate (k_d) and apparent equilibrium dissociation constant (K_b) \pm standard errors were derived from the binding curves obtained using 3 determinations at differing concentrations of the respective soluble receptors.

	k _a (1/Ms)	k _d (1/s)	K _D (M)
chicken	2.3 ± 2.6	8.8 ± 5.1 ×	5.4 ± 2.9 ×
GFRα-4 rat GFRα-4	x 10 ⁴ 2.7 ± 1.6	10 ⁻⁴ 1.1 ± 0.2 ×	10 ⁻⁹ 5.9 ± 2.8 x
	x 104	10-3	10-9

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Although apparent K_D values were very similar for both fusion proteins, R_{MAX} values were significantly different. Binding levels of ~1000 relative units (RU) were routinely obtained with chicken GFR α -4-IgGFc, whereas binding levels of rat GFR α 4-IgGFc were approximately 20 times lower, around 50-60 RU. This could be due to differences in the concentration of active chicken GFR α 4-IgGFc and rat GFR α 4-IgGFc fusion protein. The calculated equilibrium dissociation constant K_D of 5.9 \pm 2.8 nM (n=3) suggests that rat GFR α 4-1 is a receptor specific for persephin.

Northern blot analysis.

Northern blots containing 2 µg of poly(A)-rich RNA 20 derived from different rodent tissues (mouse $\mathtt{MTN}^{\mathtt{TM}}$ blot, mouse embryo MTN $^{\text{TM}}$ blot and rat MTN $^{\text{TM}}$ blot; Clontech Laboratories) were hybridized according to the manufacturer's instructions with a $\alpha\text{-}[^{32}P]\text{-}dCTP$ random-priming labeled (HighPrime kit, Roche 25 Diagnostics) 948 bp fragment derived from the rat $GFR\alpha-4$ coding sequence (as in sequence ID No. 6). Stringency washes were performed in 0.1x SSC / 0.1 %SDS at $50\,^{\circ}\text{C}$ (the two mouse blots) or $55\,^{\circ}\text{C}$ (the rat blot). The results are shown in Figure 4. In rat, a 30 very weak signal could be detected around 2.3 kb in heart, brain, liver and testis. An even weaker second transcript was present around 1.4 kb in the same tissues. In mouse, a 1.35 kb transcript was most

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intense in brain and testis, with a much weaker signal present around 2 kb. Very low mRNA expression of the 1.35 kb transcript was also present in 15-day and 17-day mouse embryo. The size of the smaller transcript is in agreement with the predicted size of the GFR α -4 coding sequence (± 880 bp + the 3' untranslated region of 570 bp).

<u>Chromosomal localization of rat, mouse and human GFRα-4.</u>

A 0.95 kb rat $GFR\alpha-4$ cDNA fragment (containing the full rat GFR α -4 coding sequence of variant A) and a 2.3 kb fragment corresponding to the rat $GFR\alpha-4$ genomic sequence (see sequence ID No. 5) were used as 15 probes in fluorescent in situ hybridization (FISH) analysis on rat chromosomes. These probes were partially overlapping. A mixture of the two probes (1 µg total DNA) was labeled with digoxigenin-11-dUTP (Roche Diagnostics, Mannheim, Germany) by nick 20 translation (Life Technologies) producing a final probe fragment size of 200-400 bp. The labeled probe was mixed with hybridization buffer (50% formamide, 2x SSC, 10% dextran sulfate). After denaturation, the mixture was placed on metaphase rat chromosome slides 25 (Islam and Levan, 1987, Helou et al., 1998) denatured at 72°C for 2 min in 70% formamide, 2x SSC. After hybridization for 48 h at 37°C, preparations were washed for 15 min in 50 % formamide, 2x SSC. Detection of labeled chromosomes was done by standard FITC anti-30 digoxigenin. Chromosome spreads were counterstained with 4', 6-diamidino-2-phenylindole (DAPI). Results were derived from micrographs of 100 different cells. Due to the small size of the probes there was

considerable background. However, most studied cells showed label at position RNO3q36 (Figure 5). About 35 % of the metaphase studies showed "double spot" label at both homologues of RNO3, whereas about 50 % had double spots on only one of the homologues or "single 5 spot" label on both homologues. No other chromosomal site showed label in several cells. Based on comparative mapping, the corresponding mouse locus would be expected to be located at MMU2 (band F), whereas a possible human location for the gene would 10 be HSA2, HSA15 or HSA20. In agreement with this, the genomic mouse and human $\mbox{GFR}\alpha\mbox{-4}$ sequences identified in the EMBL database (Accession No. AF155960 for mouse and AC017113 for human) are derived from mouse chromosome 2 (BAC clone 389B9) and from human 15 chromosome 2 (EMBL accession number AC013324; BAC388 K 24map2), respectively.

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List of abbreviations

	ARTN	artemin
	AKIN	
	BLAST	basic local alignment search tool
5	bp	base pairs
	cDNA	complementary DNA
	CNS	central nervous system
	EST	expressed sequence tag
	EVN	enovin
10	GDNF	glial cell-line derived neurotrophic
		factor
	GFRα	GDNF family receptor α
	GPI	glycosyl phosphatidyl inositol
	NTN	neurturin
15	PCR	polymerase chain reaction
	PNS	peripheral nervous system
	PSP	persephin
	SSP	smallest sum probability
	TGF-β	transforming growth factor β
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Claims

- 1. An isolated or substantially pure form of a nucleic acid molecule encoding a mammalian GDNF family receptor α -4 (GFR α -4).
 - 2. The nucleic acid molecule of claim 1 which is derived from a rat, mouse or human.
- 10 3. The nucleic acid molecule of claim 1 or 2 encoding a mammalian GDNF family receptor α -4 (GFR α -4) having the amino acid sequence illustrated in Sequence ID No. 8 or 9 or encoding a functional equivalent or bioprecursor of said receptor.

 A nucleic acid molecule according to any of claims 1 to 3 which is a DNA molecule.

- A nucleic acid molecule according to claim
 4, wherein said DNA molecule is a cDNA molecule.
 - 6. A nucleic acid molecule according to any preceding claim having the sequence illustrated in any of SEQ ID Nos 5, 6, or 7 or the complementary sequence thereof.
 - 7. A nucleic acid molecule capable of hybridising to the molecule of any of claims 1 to 6 or the complementary sequences thereof under conditions of high stringency.
 - 8. A GFR α -4 receptor encoded by a nucleic acid molecule according to any of claims 1 to 6.

- 9. A DNA expression vector comprising a nucleic acid molecule according to any of claims 4 to 6.
- 10. A host cell transformed or transfected with 5 the vector according to claim 9.
 - 11. A host cell according to claim 10, which cell is a eukaryotic cell.
- 10 12. A host cell according to claim 10 or 11 wherein said cell is a mammalian cell.
- 13. A host cell according to claim 12 which cell is a human embryonic kidney cell HEK293 or a Cos-7 cell.
 - 14. A transgenic cell, tissue or organism comprising a transgene capable of expressing a GFRα-4 receptor protein having the amino acid sequence illustrated in Sequence ID No's. 8 or 9 or the amino acid sequence of a functional equivalent or bioprecursor thereof.
- 15. A transgenic cell tissue or organism 25 according to claim 14, wherein said transgene comprises a nucleic acid molecule according to any of claims 1 to 6.
- 16. A GFRα-4 receptor protein or a functional 30 equivalent derivative or bioprecursor thereof, expressed by the cell according to any of claims 10 to 15.

- 17. A HEK293 or Cos-7 cell line trasfected or transformed with the expression vector of claim 9.
- 18. An antisense molecule comprising a nucleic acid which is capable of hybridising to the nucleic acid according to any of claims 1 to 6.
 - 19. A molecule according to claim 18 for use as a medicament.

- 20. Use of a molecule according to claim 18 in the manufacture of a medicament for treating pain or carcinoma.
- 15 21. An isolated receptor having the amino acid sequence as illustrated in any of SEQUENCE ID No 8 or 9 or the amino acid sequence of a functional equivalent or bioprecursor of said receptor.
- 20 22. A pharmaceutical composition comprising a nucleic acid molecule according to any of claims 1 to 6 together with a pharmaceutically acceptable carrier, diluent or excipient therefor.
- 25 23. A pharmaceutical composition comprising a molecule according to claim 18 or a receptor according to claim 21 together with a pharmaceutically acceptable carrier, diluent or excipient therefor.
- 30 24. A compound which acts as an agonist or an antagonist in relation to the receptor of claim 21.
 - 25. A pharmaceutical composition comprising an agonist or an antagonist according to claim 24

together with a pharmaceutically acceptable carrier, diluent, or excipient therefor.

- 26. A method of determining whether a compound is an agonist or an antagonist in relation to a receptor GFRα-4 according to any of claims 8 or 21, which method comprises contacting a cell expressing said receptor with said compound to be tested and monitoring the level of any GFRα4 mediated functional or biological response.
 - 27. A method according to claim 26 wherein said cell is a cell according to any of claims 10 to 15.
- 15 28. A method according to claim 26 or 27 wherein the GFRα-4 mediated functional or biological response comprises the level of phosphorylation in said cell.
- 29. A method of determining whether a compound is an agonist, antagonist or a ligand in relation to GFRα-4 receptor, according to claims 8 or 11, which method comprises contacting a membrane preparation of cells expressing said GFRα-4 with said compound in the presence of cRET or similar protein which interacts with GFRα-4 in the signal transduction pathway of which GFRα4 is a component and monitoring the level of any interaction of GFRα-4 with cRET or said similar protein.
- 30 30. A method of producing an antagonist or agonist of $GFR\alpha-4$ comprising the steps of a method of any one of claims 26 to 29; and additionally
 - (i) synthesizing the compound obtained or identified in said method or a

physiologically acceptable analog or derivative thereof in an amount sufficient to provide said antagonist or agonist in a therapeutically effective amount to a patient; and/or

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(ii) combining the compound obtained or identified in said method or an analog or derivative thereof with a pharmaceutically acceptable carrier.

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A compound identifiable as an agonist by the method according to any of claims 26 to 29 for use as a medicament.

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Use of a compound identifiable as an agonist by the method according to any of claims 26 to 29 in the preparation of a medicament for the treatment of neurodegenerative diseases, Alzheimers disease, Parkinsons disease, Motor Neuron Disease, peripheral neuropathy, spinal cord injury, familial hirschsprung disease, carcinomas and diseases associated with $\mbox{GFR}\alpha 4$ receptor dysfunction.

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33. A compound identifiable as an antagonist by the method according to any of claims 26 to 29 for use as a medicament.

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Use of a compound identifiable as an antagonist by the method according to any of claims 26 to 29 in the preparation of a medicament for the treatment of carcinomas or in alleviating pain.

A pharmaceutical composition comprising a compound according to claim 31 or 32 together with a

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pharmaceutically acceptable carrier, diluent or excipient therefor.

- 36. An antibody specific for $GFR\alpha-4$ receptor protein having an amino acid sequence as illustrated in Sequence ID No's. 8 or 9 or an amino acid sequence of a functional equivalent or bioprecursor of said receptor.
- 37. A pharmaceutical composition comprising an antibody according to claim 36 together with a pharmaceutically acceptable carrier, diluent or excipient therefor.
- 15 38. A method of identifying ligands for $GFR\alpha-4$ receptor protein, which method comprises contacting a receptor according to claim 8 or 11 with a cell extract or a compound to be tested and isolating any molecules bound to said receptor.
 - 39. A method of determining whether a compound is a ligand for $GFR\alpha-4$ receptor, which method comprises contacting a cell expressing said receptor according to any of claims 10 to 15 with said compound and monitoring the level of any $GFR\alpha-4$ mediated functional or biological response.
 - 40. A method according to claim 39 which comprises monitoring the level of phosphorylation in said cell.
 - 41. A compound identifiable as a ligand for $GFR\alpha-4$ according to the method of claims 39 or 40 for use as a medicament.

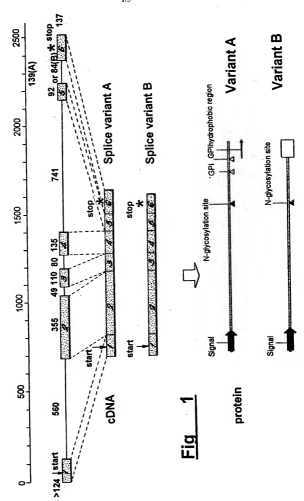
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- 42. Use of a compound identifiable according to the method of claims 39 or 40 in the preparation of a medicament for the treatment of neurodegenerative diseases, Alzheimers disease, Parkinsons disease, Motor Neuron Disease, peripheral neuropathy, spinal cord injury, familial hirschsprung disease in addition to carcinoma and diseases associated with GFR α -4 dysfunction.
- 43. A kit for determining whether a compound is an agonist or an antagonist of GFRα-4 receptor protein which kit comprises a cell according to any of claims 10 to 15, means for contacting said cell with said compound and means for monitoring the level of GFRα-4 mediated functional or biological response in said cell.
 - 44. A kit according to claim 43, wherein said $GFR\alpha-4$ mediated functional or biological response comprises the level of phosphorylation in said cell.
 - 45. A diagnostic kit including a probe which comprises any of, a nucleic acid molecule according to any of claims 1 to 6 or a fragment thereof or an antisense molecule according to claim 18 and means for contacting biological material to be tested with said probe.
- 46. A kit for determining whether a compound is a ligand of GFRα-4 receptor protein, which kit comprises a membrane preparation from cells expressing GFRα-4, means for contacting said preparation with said compound in the presence of cRET or a similar protein involved in the signal transduction pathway of

which GFR α -4 is a component and means for measuring any interaction between GFR α -4 and cRET or said similar protein.

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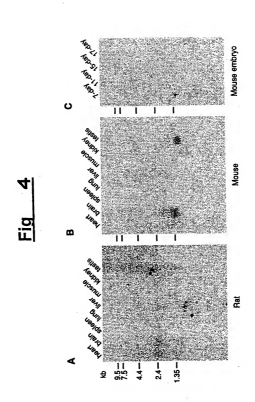
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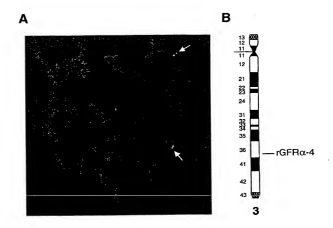
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Fig 5



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Arg Pro Arg Leu Phe Ala Phe Gln Ala Ser Cys Ala Pro Ala Pro Gly
145 150 155 160

Ser Arg Asp Gly Cys Pro Glu Glu Gly Gly Pro Arg Cys Leu Arg Ala 165 170 175

Tyr Ala Gly Leu Val Gly Thr Val Val Thr Pro Asn Tyr Leu Asp Asn 180 185

Val Ser Ala Arg Val Ala Pro Trp Cys Gly Cys Glu Ala Ser Gly Asn

195 200 205

Arg Glu Glu Cys Glu Ala Phe Arg Lys Leu Phe Thr Arg Asn Pro 210 215 220

Cys Leu Asp Gly Ala Ile Gln Ala Phe Asp Ser Ser Gln Pro Ser Val 225 230 235 240

Leu Gln Asp Gln Trp Asn Pro Tyr Gln Asn Ala Gly Cys Cys Phe Leu 245 250 255

Trp Val Ser Ser Met Ser Ile Leu Thr Ala Leu Ala Leu Gln Ala Leu 260 265 270

Leu

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<212> PRT

<213> Rattus rattus

<400> 9

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Ala Val Leu Trp Ser Leu Gly Cys Gln Arg Gly Ser Ala Ser Ser Thr 20 25 30

Glu Gly Asn Arg Cys Val Glu Ala Ala Glu Ala Cys Thr Ala Asp Glu 35 40 45

Gln Cys Gln Gln Leu Arg Ser Glu Tyr Val Ala Gln Cys Leu Gly Arg 50 55 60

Ala Gly Trp Arg Gly Pro Gly Ser Cys Val Arg Ser Arg Cys Arg Arg 65 70 75 80

Ala Leu Arg Arg Phe Phe Ala Arg Gly Pro Pro Ala Leu Thr His Ala 85 90 95

Leu Leu Phe Cys Gly Cys Glu Gly Pro Ala Cys Ala Glu Arg Arg Arg 100 105 110

Gln Thr Phe Ala Pro Ala Cys Ala Phe Ser Gly Pro Gln Leu Ala Pro 115 120 125

Pro	Ser 130	Cys	Leu	Lys	Pro	Leu 135	Asp	Arg	Cys	Glu	Arg 140	Ser	Arg	Arg	Çys
Arg 145	Pro	Arg	Leu	Phe	Ala 150	Phe	Gln	Ala	Ser	Cys 155	Ala	Pro	Ala	Pro	Gly 160
Ser	Arg	Asp	Gly	Cys 165	Pro	Glu	Glu	Gly	Gly 170	Pro	Arg	Cys	Leu	Arg 175	Ala
Tyr	Ala	Gly	Leu 180	Val	Gly	Thr	Val	Val 185	Thr	Pro	Asn	Tyr	Leu 190	Asp	Asn
Val	Ser	Ala 195	Arg	Val	Ala	Pro	Trp 200	Cys	Gly	Cys	Glu	Ala 205	Ser	Gly	Asn
Arg	Arg 210		Glu	Cys	Glu	Ala 215	Phe	Arg	Lys	Leu	Phe 220	Thr	Arg	Asn	Pro
Cys 225		Asp	Gly	Ala	Ile 230		Ala	Phe	Asp	Ser 235	Ser	Gln	Pro	Ser	Val 240
Leu	Gln	Asp	Gln	Trp		Pro	Tyr	Glm	Asn 250	Ala	Gly	Gln	Ala	Lys 255	Val

Glu Ala

INTERNATIONAL ARCH REPORT

al Application No PCT/EP 00/04918

C1201/68

A. CLASSIFICATION OF SUBJECT MATTER IPC 7 C12N15/11 C12N15/12 G01N33/50 A61K48/00

C. DOCUMENTS CONSIDERED TO BE RELEVANT

CO7K14/71

A61K38/17

Relevant to claim No.

1,3-5,

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

X

Minimum documentation searched (classification system followed by classification symbots) IPC 7 C12N C07K C12Q A61K G01N

Category . Citation of document, with indication, where appropriate, of the relevant passages THOMPSON J ET AL.: "GFRalpha-4, a new

Documentation searched other than minimum documentation to the extent that such documents are included. In the fields searched

Electronic data base consulted during the International search (name of data base and, where practical, search terms used)

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Y	cited in the application the whole document		22,23, 26-29, 37-40, 43-46
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° Special ca	her documents are listed in the continuation of box C. ategories of cited documents: ent defining the general state of the art which is not	Patent family members are listed It ster document published after the Interpropry data and not in conflict with cited to understand the principle or in.	rnational filing date the application but
"E" earlier filling ("L" docum which citatio "O" docum other "P" docum	dered to be of particular relevance document but published on or after the international	tribution of the control of the control of the control of the control of particular relevance; the control of particular relevance of the control of the con	taimed invention be considered to cument is taken alone laimed invention ventive step when the pre other such docu- us to a person skilled
	actual completion of the international search	Date of mailing of the international se	arch report
	7 November 2000	04/12/2000	
Name and	malling ackress of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL – 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	van de Kamp, M	

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FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. Claims: 1-5,7-46 (all partially); 6 (completely)

A nucleic acid according to SEQ ID NOs 5, 6 or 7 encoding a rat GDNF family receptor alpha-4 (rGFRalpha-4) according to SEQ ID NOs 8 or 9 or encoding a functional equivalent, and said encoded GFRalpha-4 proteins. Hybridizing nucleic acid molecules, vectors, hosts, transgenic cells, tissues or organisms, and pharmaceutical compositions. Agonists, antagonists and ligands, methods for their identification and use thereof. Antibodies and their use. Kits.

2. Claims: 1-5.7-46 (all partially)

A nucleic acid encoding a mouse GDNF family receptor alpha-4 (mGFRalpha-4) which is a functional equivalent of the receptor encoded by SEQ ID NOs 8 or 9, and said encoded mGFRalpha-4 protein. Hybridizing nucleic acid molecules, vectors, hosts, transgenic cells, tissues or organisms, and pharmaceutical compositions. Agonists, antagonists and ligands, methods for their identification and use thereof. Antibodies and their use. Kits.

3. Claims: 1-5,7-46 (all partially)

A nucleic acid encoding a human GDNF family receptor alpha-4 (hGFRalpha-4) which is a functional equivalent of the receptor encoded by SEQ ID NOs 8 or 9, and said encoded hGFRalpha-4 protein. Hybridizing nucleic acid molecules, vectors, hosts, transgenic cells, tissues or organisms, and pharmaceutical compositions. Agonists, antagonists and ligands, methods for their identification and use thereof. Antibodies and their use. Kits.

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

Continuation of Box I.2

Claims Nos.: 24, 25, 31-35

Remark (2): Claims 24, 25, and 31-35 refer to agonists and/or antagonists of the polypeptide(s) without giving a true technical characterisation. Moreover, no specific compounds are defined in the application. In consequence, the scope of said claims is ambiguous and vague, and their subject-matter is not sufficiently disclosed and supported (Art. 5 and 6 PCT). No search can be carried out for such purely speculative claims whose wording is, in fact, a mere recitation of the results to be achieved.

The applicant's attention is drawn to the fact that claims, or parts of claims, relating to inventions in respect of which no international search report has been established need not be the subject of an international preliminary examination (Rule 66.1(e) PCT). The applicant is advised that the EPO policy when acting as an International Preliminary Examining Authority is normally not to carry out a preliminary examination on matter which has not been searched. This is the case irrespective of whether or not the claims are amended following receipt of the search report or during any Chapter II procedure.

INTERNATION SEARCH REPORT

Information on patent family members

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